AMENDMENTS TO THE CLAIMS

- 1. (Canceled)
- 2. (Currently amended) A method according to claim 14, further including determining, based on the alignment having the smallest error, at least one of: a mismatch number based on mismatches between the first sequence and the second sequence based on the alignment having the smallest error, and, an alignment number based on matches between the first sequence and the second sequence based on the alignment having the smallest error.
- (Original) A method according to claim 2, where: the mismatches are negative matches, and, the matches can be at least one of perfect matches and positive matches.
- (Currently Amended) <u>A method for comparing a first sequence and a second sequence, the method comprising:</u>
- (a) determining a plurality of alignments of the first sequence and the second sequence,
- (b) for each said alignment of the first sequence with the second sequence, associating an errors with the said alignments of the first sequence and the second sequence.
- (c) comparing the alignment errors to identify the alignment having the smallest error, and,
- (d) based on the alignment having the smallest error, computing: a first percent identity relative to the first sequence, and a second percent identity relative to the second sequence, and
- (e) based upon the computed first percent identity and second percent identity outputting a result to at least one of a user, a storage device, a computer, or a display,A method-according-to-claim-1;

where computing a first percent identity relative to the first sequence includes: determining an alignment number based on the matches between the first sequence and the second sequence based on the alignment having the smallest error, and, forming a ratio based on the alignment number and the length of the first sequence.

- (Original) A method according to claim 4, where: the mismatches are negative matches, and, the matches can be at least one of perfect matches and positive matches.
- 6. (Currently amended) A method according to claim 14, where computing a second percent identity relative to the second sequence includes: determining an alignment number based on the matches between the first sequence and the second sequence based on the alignment having the smallest error, and, forming a ratio based on the alignment number and the length of the second sequence.
- (Original) A method according to claim 6, where: the mismatches are negative matches, and, the matches can be at least one of perfect matches and positive matches.
- 8. (Currently amended) A method according to claim 46, further including computing a third percent identity relative to the alignment having the smallest error.
- 9. (Original) A method according to claim 8, where computing a third percent identity includes: determining an alignment number based on the matches between the first sequence and the second sequence based on the alignment having the smallest error, and, forming a ratio based on the alignment number and the length of the alignment.
- 10. (Currently Amended) A method according to claim 14, further including, determining whether at least one of the first percent identity and the second percent identity is greater than a percent identity threshold.
- (Original) A method according to claim 10, further including providing a percent identity threshold.

- 12. (Currently amended) A method according to claim 44, further including, for the alignment having the smallest error, determining a number of gaps in the first sequence, and a number of gaps in the second sequence, and determining at least one of: a number based on the gaps in the first sequence based on the alignment having the smallest error, and, a number based on the gaps in the second sequence based on the alignment having the smallest error.
- 13. (Currently amended) A method according to claim 14, further including: providing at least one database, the at least one database including at least one sequence, and, retrieving at least one of the first sequence and the second sequence from the at least one database.
- 14. (Currently amended) A method according to claim 14, where, the first sequence includes comprises at least one sequence chosen from a group comprisingof: at least one-polypeptide sequences and at least one-nucleotide sequences, and, the second sequence includes comprises at least one sequence chosen from a group comprisingof: at least one-polypeptide sequences and at least one-nucleotide sequences.
- 15. (Currently amended) A method according to claim 46, where associating errors with alignments includes, aligning the first sequence and the second sequence, and, computing an error based on the number of mismatches in the alignment.
- 16. (Currently amended) A method according to claim 415, where associating errors with alignments includes, aligning the first sequence with the second sequence based on at least one insertion event in at least one of: the first sequence and the second sequence.
- 17. (Currently amended) A method according to claim 116, where associating errors includes computing a string edit distance.
- 18. (Currently amended) A method according to claim <u>116</u>, where associating errors includes comparing a number of alignment errors to an alignment error threshold.

- 19. (Currently amended) A method according to claim 14, where associating errors with alignments includes, comparing a length of the first sequence to a length of the second sequence to identify a shorter sequence and a longer sequence, and, aligning at least the entirety of the shorter sequence with at least a fragment of the longer sequence.
- 20. (Original) A method according to claim 19, where aligning at least the entirety includes inserting at least one gap into at least one of the shorter sequence and the longer sequence.
- 21. (Original) A method according to claim 19, where comparing includes, determining that the first sequence length is equal to the second sequence length, and, associating the first sequence with the shorter sequence and the second sequence with the longer sequence, and performing the aligning, and, associating the first sequence with the longer sequence and the second sequence with the shorter sequence, and performing the aligning.
- 22. (Original) A method according to claim 19, where comparing includes, determining that the first sequence length is equal to the second sequence length, and, associating at least one of: the first sequence with the shorter sequence and the second sequence with the longer sequence, and, the first sequence with the longer sequence and the second sequence with the shorter sequence.

23. - 25. (Canceled)

- 26. (Currently amended) A method according to claim 14, further including: comparing the length of the first sequence with the length of the second sequence, and, performing the alignments based on the length comparison and a percent identity threshold.
- 27. (Currently amended) A method according to claim 14, further including providing at least one interface to perform at least one of: identify the first sequence,

identify the second sequence, provide a percent identity threshold, and provide an alignment error threshold.

- 28. (Currently amended) A method according to claim 14, further comprising outputting the first percent identity and the second percent identity.
- 29. (Currently amended) A method according to claim 14, further comprising outputting the first percent identity and the second percent identity based on at least one of: a percent identity threshold and an alignment error threshold.
- 30. (Currently amended) A method according to claim 14, further comprising outputting a scoring matrix associated with the first percent identity and the second percent identity.
- 31. (Currently amended) A method according to claim 14, further comprising outputting data based on a comparison of the first percent identity and the second percent identity with a percent identity threshold.
- 32. (Currently amended) A method according to claim 14, further comprising: iteratively, storing the first percent identity and the second percent identity, retrieving at least one of a first sequence and a second sequence, and, returning to associating errors, to provide at least one stored first percent identity and second percent identity.
- 33. (Original) A method according to claim 32, where storing includes associating the first percent identity and the second percent identity with at least one of the first sequence and the second sequence.
 - 34. 46. (Canceled)
- 47. (New) A method for comparing a first sequence and a second sequence, the method comprising:
 - (a) comparing a length of the first sequence to a length of the second sequence;

(b) based on the length comparison and a percent identity threshold, determining a plurality of alignments of the first sequence with the second sequence,

where determining an alignment comprises, based on the lengths of the first and second sequences, identifying a shorter sequence and a longer sequence and aligning the shorter sequence with at least a fragment of the longer sequence;

- (c) for each said alignment of the first sequence with the second sequence, associating an error therewith, where associating an error with an alignment includes,
- i) computing an error based on the number of mismatches in the said alignment; and
- ii) computing an error based on at least one insertion event in at least one of the shorter sequence and the fragment of the longer sequence;
- (d) comparing the associated alignment errors to identify the alignment having the smallest error, and,
- (e) based on the alignment having the smallest error, computing a first percent identity relative to the first sequence, and a second percent identity relative to the second sequence,

where computing a percent identity relative to a given sequence includes determining an alignment number based on the matches between the first sequence and the second sequence based on the alignment having the smallest error, and, forming a ratio based on the alignment number and the length of the given sequence; and

(f) based upon the computed first percent identity and second percent identity outputting a result to at least one of a user, a storage device, a computer, or a display.

- 48 (New) A method according to claim 32, further comprising: sorting the at least one stored first percent identity and second percent identity based on percent identity, and, outputting the sorted first percent identity and second percent identity.
- 49. (New) A method according to claim 4, where at least one of the first sequence and the second sequence includes an ASCII string.
- 50. (New) A method according to claim 4, further comprising: performing in at least one parallel processing thread, storing the first percent identity and the second percent identity, and, retrieving at least one of a first sequence and a second sequence, and, returning to associating errors, to provide at least one stored first percent identity and second percent identity.
- 51. (New) A method according to claim 49, where at least one of the first sequence and the second sequence is selected from a database of sequences.
- 52 (New) A method according to claim 51, where the first sequence is selected from a first database of sequences and the second sequence is selected from a second database of sequences.
- 53. (New) A method according to claim 51, where the first sequence and the second sequence are selected from the same database of sequences.
- 54 (New) A method according to claim 19, further comprising aligning at least the entirety of the shorter sequence with at least a fragment of the longer sequence based on at least one insertion event in at least one of: the entirety of the shorter sequence and at least a fragment of the longer sequence.
- 55 (New) A method according to claim 54, where associating errors with alignments includes computing an error based on the number of mismatches in the alignment.

- 56. (New) A method according to claim 54, where associating errors with alignments includes computing a string edit distance.
- 57. (New) A method according to claim 55, where associating errors with alignments includes comparing a number of alignment errors to an alignment error threshold.